

# Indian genetic disease database

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## ABSTRACT

**Indians, representing about one-sixth of the world population, consist of several thousands of endogamous groups with strong potential for excess of recessive diseases. However, no database is available on Indian population with comprehensive information on the diseases common in the country. To address this issue, we present Indian Genetic Disease Database (IGDD) release 1.0 (<http://www.igdd.iicb.res.in>)—an integrated and curated repository of growing number of mutation data on common genetic diseases afflicting the Indian populations. Currently the database covers 52 diseases with information on 5760 individuals carrying the mutant alleles of causal genes. Information on locus heterogeneity, type of mutation, clinical and biochemical data, geographical location and common mutations are furnished based on published literature. The database is currently designed to work best with Internet Explorer 8 (optimal resolution 1440 × 900) and it can be searched based on disease of interest, causal gene, type of mutation and geographical location of the patients or carriers. Provisions have been made for deposition of new data and logistics for regular updation of the database. The IGDD web portal, planned to be made freely available, contains user-friendly interfaces and is expected to be highly useful to the geneticists, clinicians, biologists and patient support groups of various genetic diseases.**

## INTRODUCTION

The load of genetic diseases varies widely between different populations depending on its structure, reproductive practices and other factors. Control and management of the genetic disorders depend on identification of the variants in the genome that are causally linked with the disease. The spectrum of such variants, i.e. mutations, is different in different population groups. Remarkable progress has been made towards capturing the genomic variation in the context of genetic diseases with the advancement of DNA sequencing technologies, the capacity to handle large amount of data by building databases and faster dissemination of information through the worldwide web. It is, therefore, not surprising that the initial modest beginning of Mendelian Inheritance of Man (MIM) transformed later to Online MIM (OMIM). Currently, the most expanded version of database specifically cataloging the mutations relating genetic diseases across globe is Human Gene Mutation Database (HGMD). In addition, special interest groups generated ‘locus specific databases’ (LSDBs) and lately ‘national and ethnic mutation databases’ (NEMDBs) have also emerged containing mutational data for specific countries (Table 1). Such endeavor enormously boosts the efforts related to diagnosis of genetic diseases, detection of carriers for disease management and control and genetic counseling to mitigate the suffering of the affected families. However, no such database on genetic diseases exists for India, a country inhabited by more than a billion people and predicted to have a high load of recessive disorders in the population.

The evolutionary history of primitive Indian ethnic groups and migration from Africa, middle-east and west Asia, southern China and south-east Asia has added to the

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**Table 1.** IGDD compared to existing<sup>a</sup> NIEMDBs (National and Ethnic Mutation Databases)

Databases	Country population (in millions)	Patients/ carriers studied	Diseases	Total mutations recorded	Unique mutations	Patient-specific records	Summary statistics provided	Launched/ last updated	Published
Finnish Disease Database (Finland)	5.30	INR	35	1362 <sup>b</sup>	INR	No	No	2002	Yes (1)
Iranian Human Mutation Database (Iran)	68.69	INR	98	466	415	No	Yes	September 2003	No
The Cypriot National Mutation Frequency Database (Cyprus)	1.05	INR	19	1478	85	No	No	August 2006	Yes (2)
The Hellenic National Mutation Database (Greece)	10.68	INR	14	3179	221	No	No	June 2006	Yes (3)
The Iranian National Mutation Frequency Database (Iran)	68.69	INR	8	2614	74	No	No	August 2006	Yes (2)
The Israeli National Genetic Database (Israel)	7.60	INR	330	2581	904	No	No	July 2010	Yes (4)
The Lebanese National Mutation Frequency Database (Lebanon)	0.02	INR	6	880	60	No	No	January 2006	Yes (5)
The Moroccan Human Mutation Database (Morocco)	28.56	INR	138	INR	229	No	No	February 2010	Yes (6)
The Serbian National Mutation Frequency Database (Serbia)	7.78	INR	6	68 <sup>c</sup>	68	No	No	April 2006	No
Thailand Human Mutation and Variation database (Thailand)	66.40	INR	119	589	518	No	Yes	August 2008	Yes (7)
Turkish Human Mutation Database (Turkey)	71.51	INR	2	57 <sup>c</sup>	57	No	No	January 2006	No
FINDbase worldwide (92 populations)	NA	INR	32	3553	1226	No	Yes	June 2009	Yes (8)
Indian Genetic Disease Database (India)	1180.16	5760	52	6647	780	Yes <sup>d</sup>	Yes	August 2010	This report

NA: Not applicable; INR: Information not retrievable.

<sup>a</sup>Currently available/accessible online; Singapore Human Mutation and Polymorphism Database is not included since the variants listed in the database are not distinctly categorized into 'mutations' or 'polymorphisms'.<sup>b</sup>Not specified whether total or unique mutations.<sup>c</sup>Database records only unique mutations.<sup>d</sup>Patient-specific record of IGDD includes personal data (e.g. age, sex, ethnicity, geographical location, etc.) and clinical and bio-chemical data.

genetic diversity of the country (9). However, religion, language and geographical location of habitat serve as barriers to random mating in the Indian population. Inbreeding is practiced in some geographical regions of India (population-inbreeding coefficient: 0.00 to 0.20) (10). Thus, the overall heterogeneity of population along with the underlying endogamy makes India, a unique case of importance with respect to a high prevalence of genetic diseases and mutations. This highlights the importance of identifying recessive diseases in the Indian groups and screening the causal genes. In addition to the overall effect of 'founder events', in some communities, the load of genetic disorder is relatively higher due to the practice of consanguineous marriage, especially in south India (11).

In March 2006, a study conducted through the March of Dimes Birth Defect Foundation, reported the birth defect prevalence in India as 64.4 (per 1000 live births) (12). Rao and Ghosh (2005) report that 1 out of 20 children admitted to hospital has a genetic disorder that ultimately account for about 1 out of 10 childhood deaths (13). In India's urban areas, congenital malformations and genetic disorders are the third most common cause of mortality in newborns (14). However, there is no common source of information to assess the load of specific genetic diseases reported in India, extent of locus and mutational heterogeneity, common mutations in the causal genes and the extent of molecular studies carried out vis-à-vis lack of it in the context of the disease load. In fact, most of the pilot studies are local and hospital based. The genetic services are also not well established and localized sporadically. The situation certainly calls for a comprehensive repository of mutational data aided by specific clinical and other relevant information of patients from different regions of India. Here we describe Indian Genetic Disease Database (IGDD), a comprehensive documentation that intends to record patient-specific mutation spectrum of genetic diseases among the Indian population that would help designing assays and diagnostic tests to detect mutations, diagnose genetic diseases and identify carriers.

## DATABASE ORGANIZATION

The logistics based on which IGDD has been created is shown schematically in Figure 1. The database offers an integrated and curated repository of experimentally characterized and reported mutations responsible for genetic disorders in Indian population. An easy-to-use web interface allows a remote user to retrieve (and submit) data through interactive web forms. The home page of IGDD provides links to other major public-domain knowledge-bases on human genetic disorders. Details of the software design, data sources, query options and other features of the database are described in the following subsections.

## Software design and implementation

The database is designed and implemented on a three-tier architecture—user/client, web-interface and RDBMS

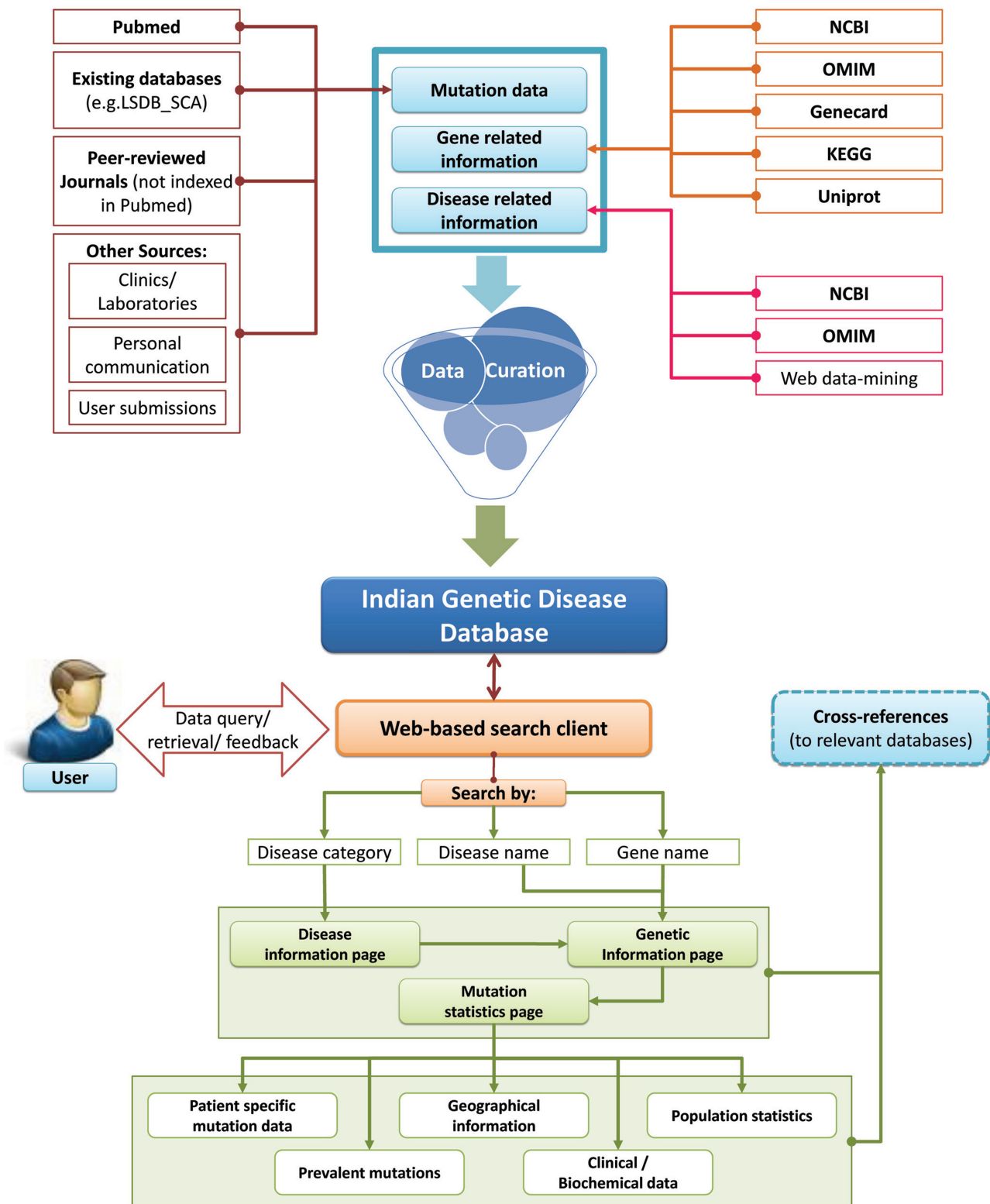


Figure 1. The schematic representation of the IGDD.

backend. The web interface is comprised of a collection of ‘web applications’/‘web forms’ developed in Microsoft Visual Basic .NET 2003. The home page of the database (<http://www.igdd.iicb.res.in>) serves as the gateway to the

interlinked web forms capable of querying the database contents dynamically as instructed (by the user) through button clicks, check-boxes and drop-down menus. In the backend, the relational database is managed with

ORACLE 9i. The data collected from different sources are initially stored in manually curated flat-files and uploaded to the database through the SQL\*Loader utility. Statistics and figures accompanying the data are auto-generated by software tools developed in-house and subject to automated revision during each update. The database is currently designed to work best with Microsoft Internet Explorer 8 (optimal resolution 1440 × 900).

### Source of data

The primary source of data is peer-reviewed published reports. With exception of a few reports all others are cited in PUBMED. In addition, data have been collected through personal communication with genetic laboratories, especially in case of β-thalassemia—the most prevalent genetic disease in India. All the data sources are duly referred to and respective bibliographic pages are hyperlinked.

For convenience of users, the diseases enlisted in IGDD have been divided into various categories such as ‘Blood Related Disorders’, ‘Eye related Disorders’, ‘Pigmentation Disorders’, etc. Diseases with complex clinical syndromes or affecting multiple organs have been included under the ‘Multisystem Disorder’ category. Every documented disorder has been described briefly and aided by proper links (to OMIM) for more detailed reading.

### Data content

IGDD release 1.0 holds entries for 52 genetic diseases and 63 related genes collated from 123 reports, published during 1993–2010. Currently, 2394 patients and 3366 carriers (resident or non-resident Indian individuals) are enlisted in the database harboring 6647 mutations of which 780 are unique in nature. Majority of these mutations are missense changes (41.3%) followed by other types of mutations (Table 2).

### Data curation

The errors found in report of mutations have been corrected when it is obvious. Those variants have not been included in the database for which coordinates of the nucleotide in the gene/cDNA and type of mutation are not clearly presented. All the mutations in the database have been linked to specific individuals with their respective phenotypic data depending on the availability of such information. Those studies that reported total mutations only, without any patient record or the number of alleles, were not enlisted in the database. Attempts are being made to convert all the mutations in single format as recommended by the Human Genome Variation Society (HGVS).

### Query options

IGDD can be navigated through by three major query options: (i) disease category, (ii) disease name and (iii) gene name, as depicted in Figure 1. Selection for a specific disease category through respective buttons directs the users to the ‘Disease Information’ page,

**Table 2.** Summary of the raw data of the IGDD

Parameters	Counts
Patients	2394
Carriers	3366
Male	920
Female	276
Sex not specified	4564
Diseases/disorders/syndromes	52
Disease with known mode of inheritance	51
Autosomal dominant	12
Autosomal recessive	29
X-linked dominant	1
X-linked recessive	6
Y-linked	0
Complex	1
Multiple	2
Genes	63
Total mutations	6647
Unique mutations	780
Missense mutations	322
Nonsense mutations	70
Deletion mutations	91
Insertion mutations	49
InDel mutations	8
Splice site mutations	48
Repeat mutations	85
Gross mutations	106
Synonymous mutations	1
Total reports studied	123
Time span (in years)	1993–2010

displaying the list of diseases under the preselected category, along with short description. Selection of a specific disease, either through the buttons in the Disease information page, or directly from a drop-down menu provided in the search bar routes the users to a ‘Genetic Information’ page that lists the causal genes, their chromosomal locations and subtypes of the disease, wherever relevant. This page may also be accessed by selecting the respective gene from a drop-down menu in the search bar. Each of the enlisted genes is linked to a ‘Mutation Statistics’ page that displays information on the encoded protein and mutation statistics along with cross references to global databases, LSDBs and Disease-Support groups.

A second level of query options is provided in the Mutation Statistics page through which the users can select for a specific type of mutation to arrive at the respective Mutation page. Figure 2 shows a screen shot of the ‘Mutation page’ that displays available individual specific-information. A search tool has been incorporated in this page to allow the user to search the relevant data for a specific mutation, either by nucleotide change or amino acid change. Moreover, a filtering utility helps the user identify mutations reported from different geographical locations of India.

The prevalent mutations for each disease gene (where  $n > 50$ ) are graphically represented in the ‘Mutation Statistics’ page. The number of individuals harboring the mutations pertaining to a specific disease from different geographical locations is pictorially represented in the Indian map. To make best of data accessibility, the

Deletion mutation Information of ATP7B																
NA: Not available, SC: Serum Caeruloplasmin, UC: Urinary Copper																
Sl.No	Donor ID	Disease Status	Age (yr.)/Sex	Nucleotide Change	Amino acid Change	Location in gene	Location in Protein	Mutation in other allele	Clinical Features	Biochemical Data	Geographical Location	Ethnic gr.	Familial / Sporadic	Remarks	Other Population	References
1	SK11*	Patient	NA/NA	c.3418delT	p.V1140A	Exon 16	ATP loop	Uncharacterized	NA	NA	North India	NA	Familial	None	Not Known	Kumar et al., 2005
2	19	Patient	10/F	c.3895delC	p.L1299L	Exon 18	ATP Hinge	Uncharacterized	Hepatic/Neurological	NA	NA	NA	Familial	None	Not Known	Santhosh et al., 2006
3	W103	Patient	8/M	c.448del5	p.E150H	Exon 2	2nd copper-binding domain	p.E150H	Hepatic/Neurological	SC: 2.5; UC: 110	East India	NA	Familial	Truncated protein with only 1st Cu binding domain	Not Known	Gupta et al., 2005
4	W166	Patient	11/M	c.892delC	p.Q298K	Exon 2	3rd copper-binding domain	p.C271X	Hepatic/Neurological	SC: 5.0; UC: 200	East India	NA	Familial	Truncated protein with only 1st 3 Cu binding domain	Not Known	Gupta et al., 2005
5	W22	Patient	18/F	c.448del5	p.E150H	Exon 2	1st copper-binding domain	Uncharacterized	Hepatic/Neurological	SC: 3.0; UC: 250	East India	NA	Familial	Truncated protein with only 1st Cu binding domain	Not Known	Gupta et al., 2005
6	W257	Patient	9/F	c.448del5	p.E150H	Exon 2	2nd copper-binding domain	p.E150H	Neurological	SC: 7.0; UC: 128	East India	NA	Familial	Truncated protein with only 1st Cu binding domain	Not Known	Gupta et al., 2005
7	W261	Patient	10/M	c.448del5	p.E150H	Exon 2	2nd copper-binding domain	Uncharacterized	Neurological	SC: 7.0	East India	NA	Familial	Truncated protein with only 1st Cu binding domain	Not Known	Gupta et al., 2007
8	W333	Patient	11/M	c.448del5	p.E150H	Exon 2	2nd copper-binding domain	p.G1061E	Hepatic/Neurological	SC: 4.0; UC: 145	East India	NA	Familial	Truncated protein with only 1st Cu binding domain	Not Known	Gupta et al., 2007
9	W378	Patient	24/M	c.448del5	p.E150H	Exon 2	3rd copper-binding	Uncharacterized	Neurological	SC: 10.4	East India	NA	Familial	Truncated protein with only 1st Cu	Not Known	Gupta et al., 2007

Figure 2. A screen-shot of the Mutation Page.

summary statistics for each disease gene has been provided as a downloadable text file (*Summary sheet*) in the Mutation Statistics page. A detailed users' manual is available in the 'Help Page' to facilitate effective usage of the database.

#### Data submission and updates

There is a provision for submission of new mutation data in the database. We shall accept both novel and previously reported mutations identified in new patients that would help project the mutational load in different population groups in India. Currently, mutation submission can be done by sending a duly filled submission form and sent via email (igdd.iicb@gmail.com). However, mutational data will be accepted based on either their publication in peer-reviewed journal or supportive documentary evidence leading to identification of the mutations. We plan to make the submission a web-based feature in near future for user convenience. All updates would be incorporated in the updated versions of the database planned to be released every 4–6 months interval depending on the volume of new data available.

#### DATABASE AVAILABILITY

The database would be publicly available free of cost without any license fees or requirement of prior registration.

#### SALIENT FEATURES OF THE DATABASE

At present, IGDD represents one of the most data-intensive repositories compared to other available NEMDBs (Table 1). It can be used as a platform to analyze and retrieve maximum information on disease prevalence trend, common mutations and most importantly the clinico-pathological data associated with specific mutations for a particular genetic disorder. In this context, unlike most other mutation databases, IGDD has been formatted as individual centric to correlate the genotype of an individual with his/her disease-related phenotype. Thus genotype–phenotype correlation could be attempted and compared between different individuals (i) who are homozygous for the same mutation or (ii) bear different mutations with similar fate of the encoded protein (e.g. different termination mutations, gross deletion, etc.). Further enrichment of the

database for this purpose would depend on the input from the investigators and we plan to make an effort toward this goal. However, since >74% of Indians inhabit in the rural areas with limited medical care and accessibility to diagnostic centers, the load of genetic diseases is expected to be much higher than projected through the database.

## CONCLUSION

Genetic diseases can be controlled best through an integrative approach of community education, population screening, genetic counseling, carrier identification and neonatal screening. IGDD would provide a key platform for clinicians, epidemiologists, geneticists and genetic counselors to access a central genetic data-source for the Indian population. This centralized mutation database is likely to play a valuable role in correlation of genotype with phenotype. We think that over long time, with enrichment of the database, the benefits accrued from it would apply to other countries (e.g. Pakistan, Bangladesh, Srilanka, Bhutan and Nepal) of the Indian subcontinent that share historically similar population groups divided by political boundaries. In addition, such implication is more directly applicable to the nonresident Indians across the world migrated in relatively recent past.

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